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(54) Title: A METHOD OF SIMULTANEOUSLY PRODUCING A LARGE NUMBER OF Leu17-VIP-ANAL AND NEW Leu¹⁷-VIP-ANALOGS

(57) Abstract

A method of simultaneously producing a large number of individual C-terminally extended analogs of Leu17-VIP is described. Said method comprises the steps of synthesizing two separate mixtures of equally long oligonucleotides coding for N-terminal and C-terminal portions of Leu17-VIP analogs, respectively, each having a few defined triplets which are made ambiguous and a few identical bases at the 3'-end, mixing said two mixtures to form a mixture of partially double stranded equally long DNA sequences, which are enzymatically converted into double stranded DNA sequences, subjecting the last mentioned mixture to cleavage, with two different enzymes to produce DNA sequences, which are inserted into similarly cleaved vectors by ligation, whereupon the vectors, are transformed into hosts, propagating said hosts to form colonies, which are analyzed one by one to establish the DNA sequences which code for a single protein, propagating separately under expression conditions those hosts which contain vectors having identified protein coding sequences, whereupon the separately expressed proteins are cleaved to release the C-terminally extended analogs of Leu17-VIP. Additionally there are disclosed new C-terminally extended analogs of Leu17-VIP and plasmids containing genes coding for the new analogs of Leu¹⁷-VIP.

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A method of simultaneously producing a large number of Leu $^{17}\text{-VIP-analogs}$ and new Leu $^{17}\text{-VIP-analogs}$.

The present invention relates to a method of simultaneously producing a large number of peptide analogs and to new peptide analogs. The invention relates specifically to a method of simultaneously producing a group of individual C-terminally extended analogs of Leu¹⁷-VIP (vasoactive intestinal polypeptide), to new C-terminally extended analogs of Leu¹⁷-VIP and to plasmids containing genes having DNA sequences which code for said new Leu¹⁷-VIP analogs.

Background

The pharmaceutical industry, and independent research workers, are constantly searching for new compounds, which slightly differ from useful known compounds, with the aim of finding more potent, more specific etc derivatives. In case the known compound of interest is a peptide or polypeptide, one or several amino acid residues of the amino acid sequence thereof are substituted for other natural or unnatural amino acid residues.

For screening purposes it would be desirable to be able to simultaneously produce a large number of analogs of a chosen peptide in order to save both time and money.

The present invention provides a method of simultaneously producing a large number of C-terminally extended analogs of Leu¹⁷-VIP, wherein all amino acid residues are of L-configuration.

Prior Art

Vasoactive intestinal polypeptide (VIP) is a

highly basic 28 amino acids long peptide with a C-terminal amide belonging to the glucagon-secretin family.

VIP was first isolated by S. Said and V. Mutt in 1970

(Said, S.I. and Mutt, V. (1970) Science 169, 1217-1218)

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from porcine upper intestinal tissue, but has later been found both in nervous tissue and in endocrine cells (Said, S.I. (1982) Vasoactive Intestinal Peptide (Raven Press, N.Y.) and (Said, S.I. (1984) Peptides 5, 143-150). The biological effects of VIP include vasodilation of cerebral blood vessels, stimulation of prolactin release, stimulation of pancreatic exocrine secretion, effect on penile erection (Said, S.I. (1982) Vasoactive Intestinal Peptide (Raven Press, N.Y.) and (Rostène, W.H. (1984) Progress in Neurobiology 22, 103-129) and Mutt, V. (1983) in Brain Peptides,

and (Rostène, W.H. (1984) Progress in Neurobiology 22, 103-129) and Mutt, V. (1983) in Brain Peptides, eds. Krieger, D.T., et al), potentiation of choliner-gically stimulated salivary flow (Lundberg, J.M., et al (1982) Acta Physiol. Scand. 114, 329-337) and acting as surfactant in the lung (Barnes, P.J. (1987) TIPS, 8, 24-27).

Recently the precursor gene coding for human VIP has been isolated and sequenced (Bodner, M., et al. (1985) Proc. Natl. Acad. Sci. USA, 82, 3548-3551) and (Linder, S., et al. (1987) Proc. Natl. Acad. Sci. 20 USA, 84, 605-609). The sequence data show that at the 3'-end of the gene coding for VIP there are three additional triplets coding for the amino acid sequence Gly-Lys-Arg. It has been shown that C-terminal Gly-Lys--Arg or Gly alone may act as substrate sites for C-ter-25 minal amidation by peptidylglycine α-amidating monooxygenase (PAMase) detected in various tissues (Bradbury, A.F., et al. (1982) Nature 298, 686-688) and (Eipper, B.A., et al (1983) Peptides 4, 921-928) and (Glembotski, C.C., et. al (1984) J. Biol. Chem. 30 259, 6385-6392) and (Gomez, S., et. al (1984) FEBS Lett. 167, 160-164) and (Eipper, B.A., et. al. (1985) Endocrinology, <u>116</u>, 2497-2504) and Ouafik, H., et. al. (1987) Proc. Natl. Acad. Sci. USA, 84, 261-264).

35 It is obvious that large scale production by genetic engineering methods of VIP, a peptide hormone of broad biological activity, and analogs thereof,

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would be desirable. Although the human precursor gene for VIP has been isolated and characterized (Bodner, M., et. al. (1985) Proc. Natl. Acad. Sci. USA, 82, 3548-3551) and Linder, S., et. al. (1987) Proc. Natl. Acad. Sci. USA, 84, 605-609), its expression in another organism has not been reported yet.

The C-terminally extended Leu¹⁷-VIP analogs of the present invention have been produced in E. coli. Description of the invention

In one aspect of the invention there is provided a method of simultaneously producing a group of individual C-terminally extended analogs of Leu¹⁷-VIP (vasoactive intestinal polypeptide), said group consisting of equally long peptides having the amino acid sequence

15 His-Ser-Asp-Ala-Val-Phe-X-Asp-Asn-Tyr-Thr-Arg-Leu-Y-1 2 3 4 5 6 7 8 9 10 11 12 13 14

-Lys-Gln-Leu-Ala-Val-Z-Lys-Tyr-Leu-Asn-Ser-Ile-Leu-20 15 16 17 18 19 20 21 22 23 24 25 26 27

-Asn-W 28 29

wherein X represents Thr, Ser, Pro or Ala, Y represents Thr, Lys, Ile or Ala Z represents Thr, Lys, Ile or Arg, and W represents Gly or Gly-Lys-Arg,

and of equally long peptides which are also $\text{Leu}^{17}\text{-VIP}\text{-}^{29}$ analogs and which arise from mutations when carrying out the method. Said method comprises the steps of

simultaneously synthesizing a first mixture of equally long oligonucleotides which during synthesis are made ambiguous at the triplets coding for the amino acid residues at positions 7 and 14 of Leu¹⁷-VIP, and which have a few identical bases at the 3'-end,

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simultaneously synthesizing a second mixture

of equally long oligonucleotides which during synthesis
are made ambiguous at the triplet coding for the amino
acid residue at position 20 of Leu¹⁷-VIP, and which
have a few identical bases at the 3'-end, these being
complementary to the few bases at the 3'-end of the
oligonucleotides in the first mixture,

mixing said first mixture with said second mixture, whereby the few bases at the 3'-end of the oligonucleotides of the first mixture anneal to the few bases at the 3'-end of the oligonucleotides of the second mixture, resulting in a mixture of partially double stranded equally long DNA sequences, which thereafter are enzymatically converted into fully double stranded DNA sequences,

subjecting said mixture of double stranded DNA sequences, which include at the 3'-end a site for cleavage with a first enzyme and at the 5'-end a site for cleavage with a second enzyme, to cleavage, simultaneously or consecutively, with the first and the second enzyme to produce DNA sequences with 3'- and 5'-ends which are ligatable to the 5'- and 3'-ends of similarly cleaved vectors,

inserting the thus cleaved DNA sequences into the thus cleaved vectors by ligation to produce a mixture of vectors, which are then transformed into hosts in per se known manner,

propagating said hosts to form colonies, which are analyzed one by one to establish those DNA sequences which code for a single protein,

propagating separately under expression conditions those hosts which contain vectors having identified protein coding sequences, the C-terminal portions of which correspond to an individual $\text{Leu}^{17}\text{-VIP-W}^{29}$ analog, whereupon the separately expressed proteins are cleaved to release the C-terminal $\text{Leu}^{17}\text{-VIP-W}^{29}$

analogs forming a group of said individual equally long peptides.

In one embodiment of this aspect of the invention, wherein W represents Gly, the $\text{Leu}^{17}\text{-VIP-W}^{29}$ analogs which arise from mutations when carrying out the method are

 $Asp^{1}-Ala^{7}-Ile^{14}-Thr^{20}-VIP-Gly^{29}$ Tyr¹-Pro⁷-Thr¹⁴-Lys²⁰-VIP-Gly²⁹ Tyr²-Thr⁷-Tyr⁸-Thr¹⁴-Thr²⁰-VIP-Gly²⁹ 10 Gly³-Thr⁷-Thr¹⁴-Arg²⁰-VIP-Gly²⁹ Pro⁴-Thr⁷-Arg¹⁴-Thr²⁰-VIP-Gly²⁹ Phe⁵-Thr⁷-Thr¹⁴-Lys²⁰-VIP-Gly²⁹ Thr⁷-Asn⁸-Arg¹⁴-Arg²⁰-VIP-Gly²⁹ Pro⁷-Gly⁸-Thr¹⁴-Lys²⁰-VIP-Gly²⁹ 15 Pro7-Gln8-Ile14-Phe19-Arg20-VIP-Gly29 Ser⁷-Pro⁸-Ile¹⁴-Arg²⁰-VIP-Gly²⁹ Ala⁷-Pro¹¹-Lys¹⁴-Lys²⁰-VIP-Gly²⁹ Pro⁷-Pro¹²-Thr¹⁴-Thr²⁰-VIP-Gly²⁹ Thr⁷-Lys¹⁴-Asn¹⁵-Ile²⁰-VIP-Gly²⁹ 20 Pro⁷-Ile¹⁴Arg¹⁵-Lys²⁰-VIP-Gly²⁹ Ala⁷-Lys¹⁴-Gln¹⁵-Arg²⁰-VIP-Gly²⁹ Ala⁷-Ile¹⁴-Thr¹⁵-Lys²⁰-VIP-Gly²⁹ Thr⁷-Lys¹⁴-His¹⁶-Lys²⁰-VIP-Gly²⁹ Pro⁷-Ile¹⁴-His¹⁶-Thr²⁰-VIP-Gly²⁹ 25 $Thr^7 - Ile^{14} - Phe^{19} - Ile^{20} - VIP - Gly^{29}$ Pro⁷-Thr¹⁴-Thr²⁰-Lys²⁴-VIP-Gly²⁹ Ser⁷-Lys¹⁴-Thr²⁰-Ile²⁴-VIP-Gly²⁹ Thr⁷-Thr¹⁴-Lys²⁰-Asp²⁸-VIP-Gly²⁹ $Gly^7 - Lys^{14} - Lys^{20} - VIP - Gly^{29}$, and $Ala^7 - Asn^{14} - Arg^{20} - VIP - Gly^{29}$. 30

In another embodiment of this aspect of the invention, wherein W represents Gly-Lys-Arg, the Leu 17 -VIP-W 29 analogs which arise from mutations when carrying out the method are

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 $Phe^{2}-Pro^{7}-Thr^{14}-Arg^{20}-VIP-Gly^{29}-Lys^{30}-Arg^{31}$ Val³-Thr⁷-Lys¹⁴-Ile²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁴-Thr¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Phe⁵-Pro⁷-Thr¹⁴-Ile²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ $\operatorname{Thr}^6 - \operatorname{Ala}^7 - \operatorname{Thr}^{14} - \operatorname{Thr}^{20} - \operatorname{VIP-Gly}^{29} - \operatorname{Lys}^{30} - \operatorname{Arg}^{31}$ 5 Thr 7-Val 8-Thr 14-Thr 20-VIP-Gly 29-Lys 30-Arg 31 Pro⁷-Pro¹¹-Thr¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹¹-Ile¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Thr 7-Arg 13-Lys 14-Arg 20-VIP-Gly 29-Lys 30-Arg 31 'Pro⁷-Gln¹³-Lys¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ 10 Pro⁷-Pro¹³-Thr¹⁴-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹³-Thr¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser⁷-Arg¹³-Thr¹⁴-Gln¹⁵-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Lys¹⁴-Gln¹⁵-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser⁷-Lys¹⁴-Asn¹⁵-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ 15 Thr 7-Lys 14-Asp 19-Lys 20-VIP-Gly 29-Lys 30-Arg 31 Thr7-Lys14-Gly19-Thr20-VIP-Gly29-Lys30-Arg31 Thr⁷-Thr¹⁴-Ile²⁰-Asp²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ala⁷-Ile¹⁴-Arg²⁰-Asp²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser⁷-Ile¹⁴-Thr²⁰-Lys²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ 20 $Pro^7 - Ile^{14} - Thr^{20} - Tyr^{25} - VIP - Gly^{29} - Lys^{30} - Arg^{31}$ and Ser⁷-Thr¹⁴-Ile²⁰-Val²⁶-VIP-Gly²⁹-Lys³⁰-Arg³¹.

Examples of hosts to be used in the above method are bacteria, e.g. of genus Bacillus or Escherichia coli, and yeasts, e.g. Saccharomyces cerevisiae. Examples of vectors to be used in the above method are bacteriophages and plasmids.

In a preferred embodiment of this aspect of the invention, the vectors are plasmids and the hosts are E. coli.

As a result of carrying out said preferred embodiment of the method of the invention there was produced a large number of C-terminally extended Leu¹⁷-VIP analogs, which form an additional aspect of the invention.

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Thus, in an additional aspect of the invention there is provided a C-terminally extended analog of Leu¹⁷-VIP which is chosen from the group consisting of peptides having the amino acid sequence

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His-Ser-Asp-Ala-Val-Phe-X-Asp-Asn-Tyr-Thr-Arg-Leu-Y-1 2 3 4 5 6 7 8 9 10 11 12 13 14

-Lys-Gln-Leu-Ala-Val-Z-Lys-Tyr-Leu-Asn-Ser-Ile-Leu-10 15 16 17 18 19 20 21 22 23 24 25 26 27

-Asn-W 28 29

wherein X represents Thr, Ser, Pro or Ala,
 Y represents Thr, Lys, Ile or Ala
 Z represents Thr, Lys, Ile or Arg, and
 W represents Gly or Gly-Lys-Arg,
 (which resulted from the method and which were originally planned to be produced)
 and

analogs of Leu¹⁷-VIP having the same amino acid sequence wherein W represents Gly,

except for His in position 1 which is substituted by Asp when X = Ala, Y = Ile and Z = Thr,

except for His in position l which is substituted by Tyr when X = Pro, Y = Thr and Z = Lys,

except for Ser in position 2 which is substituted by Tyr and Asp in position 8 which is substituted

30 by Tyr when X = Thr, Y = Thr and Z = Thr,

except for Asp in position 3 which is substituted by Gly when X = Thr, Y = Thr and Z = Arg,

except for Ala in position 4 which is substituted by Pro when X = Thr, Y = Arg and Z = Thr,

except for Val in position 5 which is substituted by Phe when X = Thr, Y = Thr and Z = Lys,

except for Asp in position 8 which is substituted by Asn when X = Thr, Y = Arg and Z = Arg, except for Asp in position 8 which is substituted by Gly when X = Pro, Y = Thr and X = Lys, except for Asp in position 8 which is substituted 5 by Gln and Val in position 19 which is substituted by Phe when X = Pro, Y = Ile and Z = Arg,except for Asp in position 8 which is substituted by Pro when X = Ser, Y = Ile and Z = Arq. except for Thr in position 11 which is substituted 10 by Pro when X = Ala, Y = Lys and Z = Lys, except for Arg in position 12 which is substituted by Pro when X = Pro, Y = Thr and Z = Thr,except for Lys in position 15 which is substituted by Asn when X = Thr, Y = Lys and Z = Ile, 15 except for Lys in position 15 which is substituted by Arg when X = Pro, Y = Ile and Z = Lys,except for Lys in position 15 which is substituted by Gln when X = Ala, Y = Lys and Z = Arg,except for Lys in position 15 which is substituted 20 by Thr when X = Ala, Y = Ile and Z = Lys, except for Gln in position 16 which is substituted by His when X = Thr, Y = Lys and Z = Lys, except for Gln in position 16 which is substituted by His when X = Pro, Y = Ile and Z = Thr,25 except for Val in position 19 which is substituted by Phe when X = Thr, Y = Ile and Z = Ile, except for Asn in position 24 which is substituted by Lys when X = Pro, Y = Thr and Z = Thr,except for Asn in position 24 which is substituted 30 by Ile when X = Ser, Y = Lys and Z = Thr, except for Asn in position 28 which is substituted by Asp when X = Thr, Y = Arg and Z = Lys, and X = Gly when Y = Lys and Z = Lys, and X = Ala when Y = Asn and Z = Arg, 35 [these analogs can be named with regard to VIP as follows:

 Asp^{1} -Ala⁷-Ile¹⁴-Thr²⁰-VIP-Gly²⁹ $_{\text{Tyr}}^{1}$ -Pro $_{\text{-Thr}}^{14}$ -Lys $_{\text{-VIP-Gly}}^{29}$ Tyr²-Thr⁷-Tyr⁸-Thr¹⁴-Thr²⁰-VIP-Gly²⁹ Gly³-Thr⁷-Thr¹⁴-Arg²⁰-VIP-Gly²⁹ Pro⁴-Thr⁷-Arg¹⁴-Thr²⁰-VIP-Gly²⁹ 5 $Phe^5-Thr^7-Thr^{14}-Lys^{20}-VIP-Gly^{29}$ Thr7-Asn8-Arg14-Arg20-VIP-Gly29 Pro⁷-Gly⁸-Thr¹⁴-Lys²⁰-VIP-Gly²⁹ Pro⁷-Gln⁸-Ile¹⁴-Phe¹⁹-Arg²⁰-VIP-Gly²⁹ Ser⁷-Pro⁸-Ile¹⁴-Arg²⁰-VIP-Gly²⁹ 10 Ala⁷-Pro¹¹-Lys¹⁴-Lys²⁰-VIP-Gly²⁹ Pro7-Pro¹²-Thr¹⁴-Thr²⁰-VIP-Gly²⁹ $Thr^7 - Lys^{14} - Asn^{15} - Ile^{20} - VIP - Gly^{29}$ Pro⁷-Ile¹⁴Arg¹⁵-Lys²⁰-VIP-Gly²⁹ Ala⁷-Lys¹⁴-Gln¹⁵-Arg²⁰-VIP-Gly²⁹ 15 Ala⁷-Ile¹⁴-Thr¹⁵-Lys²⁰-VIP-Gly²⁹ Thr⁷-Lys¹⁴-His¹⁶-Lys²⁰-VIP-Gly²⁹ Pro⁷-Ile¹⁴-His¹⁶-Thr²⁰-VIP-Gly²⁹ Thr⁷-Ile¹⁴-Phe¹⁹-Ile²⁰-VIP-Gly²⁹ Pro⁷-Thr¹⁴-Thr²⁰-Lys²⁴-VIP-Gly²⁹ ₹ 20 Ser⁷-Lys¹⁴-Thr²⁰-Ile²⁴-VIP-Gly²⁹ Thr⁷-Thr¹⁴-Lys²⁰-Asp²⁸-VIP-Gly²⁹ $Gly^7 - Lys^{14} - Lys^{20} - VIP - Gly^{29}$ and Ala⁷-Asn¹⁴-Arg²⁰-VIP-Gly²⁹] (which arose as mutations when carrying out one run 25 vention)

of the preferred embodiment of the method of the in-

and analogs of Leu¹⁷-VIP having the same amino acid sequence wherein W represents -Gly-Lys-Arg 30

except for Ser in position 2 which is substituted by Phe when X = Pro, Y = Thr and X = Arg,

except for Asp in position 3 which is substituted by Val when X = Thr, Y = Lys and Z = Ile,

except for Ala in position 4 which is substituted 35 by Pro when X = Thr, Y = Thr and Z = Lys,

except for Val in position 5 which is substituted by Phe when X = Pro, Y = Thr and Z = Ile, except for Phe in position 6 which is substituted by Thr when X = Ala, Y = Thr and Z = Thr, 5 except for Asp in position 8 which is substituted by Val when X = Thr, Y = Thr and Z = Thr, except for Thr in position 11 which is substituted by Pro when X = Pro, Y = Thr and Z = Lys,except for Thr in position 11 which is substituted 10 by Pro when X = Pro, Y = Ile and Z = Arg,except for Leu in position 13 which is substituted by Arg when X = Thr, Y = Lys and Z = Arg, except for Leu in position 13 which is substituted by Gln when X = Pro, Y = Lys and Z = Lys, 15 except for Leu in position 13 which is substituted by Pro when X = Pro, Y = Thr and Z = Thr,except for Leu in position 13 which is substituted by Pro when X = Pro, Y = Thr and Z = Arg;except for Leu in position 13 which is substituted 20 by Arg and Lys in position 15 which is substituted by Gln when X = Ser, Y = Thr and Z = Thr, except for Lys in position 15 which is substituted by Gln when X = Pro, Y = Lys and Z = Arg, except for Lys in position 15 which is substituted 25 by Asn when X = Ser, Y = Lys and Z = Arg,except for Val in position 19 which is substituted by Asp when X = Thr, Y = Lys and Z = Lys, except for Val in position 19 which is substituted by Gly when X = Thr, Y = Lys and Z = Thr, 30 except for Asn in position 24 which is substituted by Asp when X = Thr, Y = Thr and Z = Ile, except for Asn in position 24 which is substituted by Asp when X = Ala, Y = Ile and Z = Arg, except for Asn in position 24 which is substituted 35 by Lys when X = Ser, Y = Ile and Z = Thr, except for Ser in position 25 which is substituted by Tyr when X = Pro, Y = Ile and Z = Thr, and

except for Ile in position 26 which is substituted by Val when X = Ser, Y = Thr and Z = Ile, [these analogs can be named with regard to VIP as follows:

5 $Phe^{2}-Pro^{7}-Thr^{14}-Arg^{20}-VIP-Gly^{29}-Lys^{30}-Arg^{31}$ Val³-Thr⁷-Lys¹⁴-Ile²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro4-Thr7-Thr14-Lys20-VIP-Gly29-Lys30-Arg31 Phe 5-Pro7-Thr 14-Ile 20-VIP-Gly 29-Lys 30-Arg 31 Thr⁶-Ala⁷-Thr¹⁴-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ 10 Thr 7-Val 8-Thr 14-Thr 20-VIP-Gly 29-Lys 30-Arg 31 Pro⁷-Pro¹¹-Thr¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹¹-Ile¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Thr 7-Arg 13-Lys 14-Arg 20-VIP-Gly 29-Lys 30-Arg 31 Pro⁷-Gln¹³-Lys¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ 15 Pro⁷-Pro¹³-Thr¹⁴-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹³-Thr¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser7-Arg13-Thr14-Gln15-Thr20-VIP-Gly29-Lys30-Arg31 Pro7-Lys14-Gln15-Arg20-VIP-Gly29-Lys30-Arg31 Ser⁷-Lys¹⁴-Asn¹⁵-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Thr7-Lys14-Asp19-Lys20-VIP-Gly29-Lys30-Arg31 Thr7-Lys14-Gly19-Thr20-VIP-Gly29-Lys30-Arg31 Thr 7-Thr 14-Ile 20-Asp 24-VIP-Gly 29-Lys 30-Arg 31 Ala⁷-Ile¹⁴-Arg²⁰-Asp²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser⁷-Ile¹⁴-Thr²⁰-Lys²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ 25 . $Pro^7 - Ile^{14} - Thr^{20} - Tyr^{25} - VIP - Gly^{29} - Lys^{30} - Arg^{31}$ and Ser⁷-Thr¹⁴-Ile²⁰-Val²⁶-VIP-Gly²⁹-Lys³⁰-Arg³¹]

(which arose as mutations when carrying out a second 30 run of the preferred embodiment of the method of the invention).

All the amino acid residues in the C-terminally extended $\text{Leu}^{17}\text{-VIP}$ analogs of the invention are of L-configuration.

In a further aspect of the invention there is provided a plasmid containing a gene having a DNA

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sequence which codes for a C-terminally extended Leu¹⁷-

All the C-terminally extended Leu¹⁷-VIP analogs of the invention have, besides similar properties to natural human VIP, some useful properties in common, namely, they are ligands at VIP receptors, they are resistant to oxidation and can be kept at acidic pH in solution (due to the replacement of methionine in position 17 of natural VIP by leucine), and, since they are C-terminally extended, they exhibit a prolonged effect both in vitro and in vivo, which is considered to be of advantage in view of the extremly short biological half-life of VIP.

Thus the C-terminally extended Leu¹⁷-VIP analogs of the invention are useful as potential active ingredients in new pharmaceuticals, as well as in cosmetics (for hydration of the skin).

The new pharmaceuticals are intended to be administered

- 20 1) for the treatment of asthma and of constriction of the upper airways in general, specially for vasodilation in the lungs,
 - 2) as pheripheric vasodilators to be locally applied for i.a. erection of penis,
- 25 3) for the treatment of VIPOMA (a VIP producing intestinal tumour which causes death of a human by loss of 5-15 liters of water per day),
 - 4) for the treatment of disorders of blood flow, blood-pressure, intestinal motility and urinary bladder,
 - for potentiation of salivation e.g. for the treatment of blocked or reduced salivation (e.g. due to administration of antipsychotic drugs or antidepressants), and
- 35 6) for the stimulation of pancreatic juice secretion.

 The composition of pharmaceuticals including
 the active ingredient will depend on the intended

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route of administration and the specific disorder to be treated. A nasal or oral spray is suitable when asthma is to be treated. Aqueous solutions or tablets for oral administration will be suitable for other indications. In all types of pharmaceuticals the active ingredient is in admixture with pharmaceutically acceptable excipients and/or diluents normally used in the field of pharmacy.

General strategy to obtain Leu¹⁷-VIP analog-coding

DNA sequences cloned in an E. coli expression vector A large number of artificial genes coding for C-terminally extended Leu¹⁷-VIP analogs were planned to be obtained directly as parts of an E. coli plasmid expression vector by performing only one cloning step. The synthetic strategy utilized the principle of a method described originally [Simoncsits, A., Kálmán, M., Cserpán, J. and Kari, C. (1984) Nucleic Acids Res. Symp. Ser. 14, 321] for obtaining a large number of artificial E. coli promoter derivatives. Briefly, two single-stranded oligodeoxyribonucleotides were chemically synthesized and were annealed [Itakura, K. (1982) TIBS, 7, 442] through their 3'-terminal complementary sequences. The partial duplex obtained so covered the whole Leu17-VIP coding region and contained additional terminal sequences to be used for cloning. The two oligodeoxyribonucleotides were obtained by ambiguous chemical synthesis so that the mixture of all four activated nucleotide monomers were used at certain predetermined positions (i.e. in triplets coding for the amino acid residues 7, 14 and 20 of Leu¹⁷-VIP) which were outside of the annealing region. The annealed partial duplex was converted by a mutually primed synthesis effected by DNA polymerase (Klenow

fragment) into double-stranded DNA mixture containing

35 all possible homoduplexes. These were then cleaved
with two different restriction endonucleases the cleavage sites of them being located upstream and downstream



of the Leu¹⁷-VIP coding region and the cleaved mixture was ligated with the appropriately cleaved expression vector pPEX. The ligated mixture was transformed into competent E. coli cells and the recombinants containing Leu¹⁷-VIP related coding sequences were selected by colony hybridization. The positive recombinants were further analyzed by nucleotide sequencing to identify the particular mutations obtained in the cloned Leu¹⁷--VIP coding region.

During the design of the synthetic oligodeoxyribonucleotides, it was considered that they should
code for a methionine preceding the Leu¹⁷-VIP coding
region and for a Gly (VIPa analogs) or a Gly-Lys-Arg
(VIPb-analogs) extension at the carboxy terminus.

15 Methionine was included to be able to release the
Leu¹⁷-VIP analogs from the fusion protein by CNBr
cleavage, while the carboxy terminal extensions could

serve as substrate sites for amidation catalysed by peptidylglycine α-amidating monooxygenase (PAMase)

20 [Bradburg, A.F., Finnie, M.D.A. and Smyth, D.G. (1982)

Nature 298, 686]. The positions of the mixed chemical couplings whithin the nucleotide sequences were chosen

couplings whithin the nucleotide sequences were chosen to be located in the predetermined X, Y and Z codons so that after performing the steps shown in Scheme 1, the maximum number of amino acid variations be obtained

for X, Y and Z. By choosing the first (for codon X) or the second (for codons Y and Z) base of the target codons as mixed positions, four amino acids could be obtained of each three codon variations, yielding altogether 64 possible Leu¹⁷-VIP analogs after expression.

The VIP1 oligonucleotide, shown as the upperstrand of the partial, annealed duplex in Scheme 1 contains two mixed positions (N), while the lower strand VIP2 oligonucleotide contains only one. The VIP2 oligonucleotide was prepared in two variations

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carrying information for C-terminal extensions as above (VIP2a and VIP2b oligonucleotides in Scheme 1). To obtain both VIPa and VIPb analogs, two separate experiments were carried out using VIP1 + VIP2a or VIP1 + VIP2b olignucleotides, respectively.

The double stranded DNA mixture obtained by Klenow polymerase reaction contained homoduplex molecules which were separated by cloning into an expression vector. In principle, all recombinants should contain only one type of mutant gene coding for a particular Leu¹⁷-VIP analog.

pPEX vector

pPEX is a highly efficient E. coli expression vector constructed from a rac fusion [Boros, J., Lukacsovich, T., Baliko, G. and Venetianer, P. (1986) 15 Gene 42, 97] promoter vector L α int 23 named as pPEX by us. The L α int 23 vector, which expresses under rac promoter control a fusion protein composed of 280 amino acids of E. coli β -galactosidase and part of the bacterial CAT (chloramphenicol acetyl transferase) was modified by eliminating its unique BamHI site and by replacing its CAT coding region located between ClaI and EcoRI sites with a synthetic polycloning region. The relevant region of the pPEX vector obtained so is shown in Scheme 2. The vector contains unique BamHI, KpnI, SacI, ApaI and EcoRI sites in the polycloning region and β -galactosidase gene fusions can be performed using the BamHI and any of the other unique sites. When the Leu¹⁷-VIP coding genes are 30 cloned into the BamHI and EcoRI sites of the pPEX vector, the resulting gene fusions code for 314 and 316 amino acids long β -galactosidase-VIPa and β -galactosidase-VIPb fusion proteins, respectively, of which proteins 284 amino acids are derived from the fusion partner (Scheme 1).

EXAMPLE

Preparation of the mixture of DNA regions coding for Leu¹⁷-VIP analogs

MATERIALS

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Restriction enzymes and T4 DNA ligase were obtained from New England Biolabs. T4 polynucleotide kinase and Klenow polymerase were from Boehringer, $[\gamma^{-32}P]ATP$ (>5000 Ci/mmol) and $[\alpha^{-32}P]dATP$ (800 Ci/mmol) were from Amersham, Rabbit antisera directed against BSA-VIP conjugates was provided by Dr. Per Askelöf (SBL, Stockholm, Sweden). Immunoblot assay kit containing anti-rabbit IgG-alkaline phosphatase conjugate and color development reagents BCIP/NBT was purchased from Bio-Rad. T₁ RNase was purchased from Calbiochem.

15 Oligodeoxyribonucleotides

CCGGATCCATATGCACTCTGACGCTGTTTTCNCTGACAACTACACT-CGTCTGANAAAACAGCTGGCT (VIPl oligonucleotide),

AAGAATTCAGCCGTTCAGGATAGAGTTCAGGTACTTTNTAACAGCC-AGCTGT (VIP2a oligonucleotide),

AAGAATTCAACGTTTGCCGTTCAGGATAGAGTTCAGGTACTTTNTA-ACAGCCAGCTGT (VIP2b oligonucleotide) and

CAGGGTGAAACGCAGGTCGCCAGCGGC (lac Z 27-mer primer) oligonucleotides were prepared on an automatic DNA synthesizer by the phosphoramidite chemistry (Pharmacia Gene Assembler) and were purified by electrophoresis on polyacrylamide gels containing 8 M urea. Oligonucleotides were 5'-phosphorylated with $[\gamma^{-32}P]$ ATP by using T4 polynucleotide kinase [Maniates, T., Fritsch, E.F. and Sambrook, J. (1982) Molecular Cloning: A laboratory manual, Cold Spring Harbor Laboratory, Cold Spring Harbor N.Y] when they were to be used as either hybridization probe or sequencing primer.

100 pmol of VIPl oligonucleotide was mixed with 100 pmole of either VIP2a or VIP2b oligonucleotide 35 in 50 μl of water, the mixture was kept at 90°C for 3 min and was let to cool to room temperature in approx.

1 hr period. The solution was made up to 100 µl containing 10 mM Tris-HCl, pH 8.2, 5 mM MgCl₂, 50 µM each of dCTP, dGTP and TTP, 5 μ l of $[\alpha^{-32P}]$ dATP (approx. 62.5 pmol) and 1 μ l of 5U/ μ l Klenow polymerase. After 15 min 10 µl of 1 mM dNTP mixture (containing all 5 four deoxynucleoside 5'-triphosphates) was added and the solution was incubated at room temperature for 15 min. The DNA was precipitated with the help of l μl of 10 μg/μl yeast carrier tRNA (ctRNA) by ethanol precipitation and purified by 10% acrylamide-8 M urea gel electrophoresis at 400V for 2 hrs (gel thickness 0.4 mm). The gel was radioautographed, the major radioactive band was cut out and soaked in 300 µl of 50 mM NaCl - 0.1% SDS solution at 37°C for 12 hrs. The supernatant was phenol extracted and the precipitation was carried out by adding 1 µl of 10 µg/µl ctRNA, 30 μ l of 3 M NaOAc, pH 5.2 and 900 μ l of ethanol in a liquid nitrogen bath. Centrifugation (14 000 rpm, 3 min) resulted in a radioactive pellet which was washed with ethanol, dried and dissolved in sterile **120** water.

20 pmol of the radioactive Leu¹⁷-VIP-DNA obtained as above was treated with 200 units of BamHI and EcoRI each in 200 µl of 100 mM NaCl, 50 mM Tris HCl, pH 7.5, 10 mM MgCl₂, 1 mM DTT (high-salt buffer) at 37°C 25 for 20 hrs. The reaction mixture was heated to 60°C for 10 min, phenol extracted (two times), and the DNA was recovered from the waterphase by ethanol precipitation as above (two times) followed by ethanol washing, drying and redissolving in sterile water. Analysis of the mixture by 10% acrylamide-8 M urea electrophoresis showed that only approx. 50% of the duplex was cleaved and the fraction of the DNA which had been cleaved by both enzymes was only approx. 30%. Nevertheless, the mixture was used as it was 35 for cloning into BamHI-EcoRI cleaved pPEX vector.

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Cleavage of pPEX vector with BamHI and ECORI

2 µg of pPEX 13 was treated with 10-10 units
of BamHI and EcoRI in 20 µl of high salt buffer at
37°C for 4 hrs. The reaction mixture was applied onto
an agarose gel (0.5%) and the electrophoresis was
performed in TAE buffer (40 mM Tris-acetate, 2 mM
EDTA pH 7.5) at 60V for 1 hr in the presence of ethidium bromide. The linear vector band was cut out,
electroeluted, phenol extracted twice, ethanol precipitated, washed with ethanol and dried. The pellet
was dissolved in sterile water.

Cloning of Leu17-VIP-DNA into pPEX vector

0.2 µg BamHI-EcoRI cleaved pPEX vector and approx. 5 pmol BamHI-EcoRI Leu¹⁷-VIP-DNA obtained as above were reacted in 20 µl reaction mixture containing 50 mM Tris-HCl pH 7.5, 10 mM MgCl₂, 10 mM DTT, 1 mM ATP and 80 units of T4 DNA ligase at 15°C for 12 hrs. The reaction mixture was transformed into JM101 E. coli cells [genotype (supE, thi, Δ(lac-proAB), [F', traD36, proAB, lacIqZAM15), Messing, J., Crea, R. and Seeburg, P.H. (1981) Nucleic Acids Res. 9, 309-321] using a frozen stock [Hanahan, D. in DNA cloning, Vol I., edited by Glover, D.M. IRC Press Limited (1985), pp 109-135, protocol 3]. Ampicillin resistant colonies were picked for colony hybridization [Grunstein, M. and Hogness, D.S. (1975) Proc. Natl. Acad. Sci. USA 72, 3961-3965] with ³²P-labeled VIPl oligonucleotide probe. Approx. 80-90% of the colonies were positive in this test for both VIPa and VIPb mutants. 120 colonies of both series were taken for plasmid preparation and sequencing.

Plasmid preparation and sequencing

A single colony was inoculated into 3 ml of LB-medium containing 100 µg/ml ampicillin and the culture
35 was shaken for 12-16 hrs at 37°C. Plasmid DNA was
prepared by a rapid alkaline extraction procedure
[Birnboim, H.C. and Doly, J. (1979) Nucleic Acids

Res., 7, 1513-1523]. The plasmid DNA was further treated with 5 units of T_1 RNase in 100 μl solution containing 10 mM Tris-HCl, pH 8.0, 1 mM EDTA at 37°C for 30 min followed by phenol extraction and ethanol precipitation. The pellet was taken up in 30 μl of sterile water.

Dideoxynucleotide sequencing was performed on HindIII linearized plasmid template $(0.2-0.4 \mu g)$ using $5'-^{32}P$ -phosphorylated lac Z 27-mer sequencing primer (0.25 pmol) and heat denaturation as described [Hong, G.F., (1982) Bioscience Reports $\underline{2}$, 907]. Sequencing reactions for 24 clones were performed

Expression of Leul 7-VIP analogs

at the same time.

E. coli JM101 cells transformed with pPEX Leu17-VIP analog plasmids were grown at 37°C, shaking 250 rpm, in 1 ml LB medium containing 0.1 mg/ml ampicillin. When the cell density reached 0.5 at A=600 nm the cells were induced either by addition of IPTG (final concentration 2.5 mM) and shaken at 250 rpm for 4 hours at 37°C or by addition of lactose (final concentration 1%) and shaken at 250 rpm for 20 hours. After induction the cells were collected by centrifugation at 12 000 rpm in an Eppendorf centrifuge for 1 minute, lysed at 100°C for 3 minutes in 300 μ l of lysis buffer (0.125 M Tris-HCl pH 6.8, 30% glycerol, 2% SDS, 6 M urea and 25 1 M 2-mercaptoethanol) and 30 μl of non-induced extracts and 30 µl of induced extracts were run on SDS-10% polyacrylamide gels according to Laemmli, U.K. (1970) Nature, 227, 680-685 to monitor the degree of protein expression as shown by coomassie brilliant blue staining of the gels. As estimated from the gels the amount of expressed fusion protein was more than 60% of total cell proteins.

Immunological recognition/detection of β-galactosidase-35 -Leu 17-VIP analogs

6 μl aliquot of non-induced and 6 μl of 50x diluted induced samples were run on SDS-10% polyacrylamide

gels in parallel and half of the gels were stained with coomassie brilliant blue and the other half of the gels were electroblotted onto nitrocellulose papers. The blotted nitrocellulose papers were incubated with a polyclonal rabbit serum containing antibodies directed against BSA-VIP, and the immunoblot was developed by treatments with goat anti rabbit IgG conjugated alkaline phosphatase and BCIP/NBT color development solution (Bio-Rad immuno assay kit) according to the manufacturer's instructions. A very strong immunostaining was obtained with the fusion proteins containing Leu¹⁷-VIP analogs.

l liter scale purification of expressed β-galactosidase - -Leu 17-VIP analog fusion proteins

E. coli JM101 cells carrying the pPEX-Leu¹⁷-VIP 15 analog plasmids were shaken (250 rpm) in 1 liter LB medium containing 0.1 mg/ml ampicillin at 37°C. When the absorbance at 600 nm reached 0.5 the cells were induced with lactose (1% final concentration) and 20 shaken at 250 rpm at 37°C for 20 hours. After induction the cells were harvested by centrifugation at 2000 x g, 4°C for 10 minutes, washed with 40 ml of TBS, pH 7.6, and resuspended in 30 ml TBS containing 10 µM PMSF and 0.5 mM EDTA. The cell suspension was then 25 freeze-thawed and sonicated (five 1 minute bursts) in a Branson sonifier, then centrifuged at 17000 x g, 4°C, for 30 minutes. The pellet containing the fusion protein was homogenized in 50 ml 6 M guanidium chloride containing 1% 2-mercaptoethanol according to Goeddel, 30 D.V. et. al. (1979) PNAS (USA), 76, 106-110 and centrifuged at 60 000 x g for 60 minutes. The clear, amber supernatant containing the fusion protein was diluted to 100 ml with 6 M quanidium chloride - 1% 2-mercapto-

to 100 ml with 6 M guanidium chloride - 1% 2-mercaptoethanol and 172 ml TBS was added dropwise with vigorous 35 stirring which rendered most of the fusion protein to precipitate while most of the host proteins remained in solution. The slurry was stirred at room temperatur

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for 30 minutes before centrifugation at 10 000 x g for 30 minutes at 10°C . The pellet so obtained contains most of the 3-galactosidase-Leu¹⁷-VIP analog fusion protein.

CNBr_cleavage_of_fusion_protein

The partially purified β-galactosidase - Leu¹⁷-VIP analog fusion protein was dissolved by homogenization in 25 ml 70% HCOOH and 1.5 g CNBr was added per gram precipitate. The solution was stirred at room temperature for 70 hours and then evaporated at room temperature under reduced pressure using a Rotavapor.

20 ml 50 mM HCl in methanol was added per gram fusion protein to the residue and stirred at room temperature over night. Then the slurry was centrifuged at 2000 x g, 4°C, for 10 minutes and the supernatant collected.

Purification of Leu¹⁷-VIP analogs

The supernatant containing the Leu¹⁷-VIP analog was either 1)

- freeze-dried and the Leu¹⁷-VIP analog-containing
 residue was dissolved in 1% TFA and purified
 on HPLC (C₁₈-Bondapack, reverse phase)
 or 2) subjected to further purification as follows
 before the HPLC step.
- To the acidic methanol phase containing the Leu¹⁷25 -VIP analog solid NaCl (0.2 g/ml) and 4 volumes
 of ice cold diethylether were added.

The precipitate (containing Leu¹⁷-VIP analog) was collected on a sintered glass funnel (pore size no 3) and dissolved by washing with 5 ml 1 M acetic acid. The dissolved Leu¹⁷-VIP analog was desalted with a Sephadex G25 column (7 cm long, 2 cm diameter) and freeze-dried before dissolution in 1% TFA and purified on HPLC with a 20%-40% acetonitrile linear gradient in 1% TFA. The Leu¹⁷-VIP analog was eluted at 33% acetonitrile. The purified Leu¹⁷-VIP analog was then tested in biochemical and biological assay systems.

EXAMPLE: VIPa (X=Thr, Y=Arg, Z=Lys, W=Gly)

The fusion protein β-galactosidase - VIPa was expressed in 1 liter LB medium containing 0.1 mg/ml ampicillin and purified as described above. The fusion protein was cleaved with CNBr and VIPa so obtained was further purified as described above and tested in the following assay systems:

- 1) Radioimmunoassay (RIA)
- 10 2) Radioreceptor assay
 - 3) Stimulation of 3',5'-cyclic AMP production
 - 4) Bicarbonate secretion in cat in vivo

1) Radioimmunoassay (RIA)

In the VIP - RIA kit purchased from Penninsula Laboratories (USA) recombinant VIPa is recognized as VIP when measured according to the manufacturer's recommendation, based on the method of Fahrenkrug, J. and Schaffalitzky De Muckadell, O. (1977) J. Lab.

20 Clin. Med. 89, 1379-1388.

2) Radioreceptor assay

Radioreceptor assay was carried out on membranes from rat cerebral cortex using chloramin-T iodinated ¹²⁵I-VIP (Halldén, G. et. al. (1986) Reg. Pep. <u>16</u>,

- 25 183-188) at 0.5 nM concentration, studying the displacement of the labeled VIP by purified VIP from porcine intestine and by recombinant VIPa from E. coli as described by Abens, J. et al. (1984) Peptides 5, 375-377. The affinity of recombinant VIPa was within
- the experimental error identical to that of VIP purified from porcine intestine with a Kd value of 1.5-1.8 nM at equilibrium, proving that recombinant VIPa is a high affinity ligand at VIP receptors, binding in a manner indistinguishable from that of VIP obtained
- 35 from other sources.

3) Stimulation of 3',5'-cyclic AMP synthesis

Stimulation of 3',5'-cyclic AMP (cAMP) synthesis
in tissue slices from rat cerebral cortex by VIP purified from porcine and recombinant VIPa was studied
to establish whether or not recombinant VIPa, a ligand
of the VIP receptor according to the result in 2),
behaves as an agonist or an antagonist. The experiments
were carried out as follows:

Rat cerebral cortical slices (0.4 mm x 0.4 mm) were preincubated in Krebs Ringers bicarbonate buffer, bubbled with O_2/CO_2 (95%/5%) (V/V) for 60 minutes at 36°C in the presence of 10 mM theophylline (a phosphodiesterase inhibitor). The tissue slices were then incubated in 600 μ l total volume with porcine VIP at 10, 50, 100 nM, 1 and 10 μM concentration for 10 minutes and with recombinant VIPa at 10, 50, 100 and 300 nM concentration (estimated by 1)) for 10 minutes. The incubations were terminated by addition of 150 μl EGTA (100 mM) solution followed by placing the test tubes into a boiling water bath for 3 minutes. The 20 cAMP content was measured according to Brown, B.L. et al. (1972) Adv. Cycl. Nucl. Res. $\underline{2}$, 25-40. The results indicate that at 10 minutes incubation time recombinant VIPa produced an increase in cAMP levels and thus act as agonist at the VIP-receptor coupled 25 adenylate cyclase, which is thought to mediate the physiological actions of VIP (cf. Rostène, W.H. (1984) Progr. Neurobiol. <u>22</u>, 103-129).

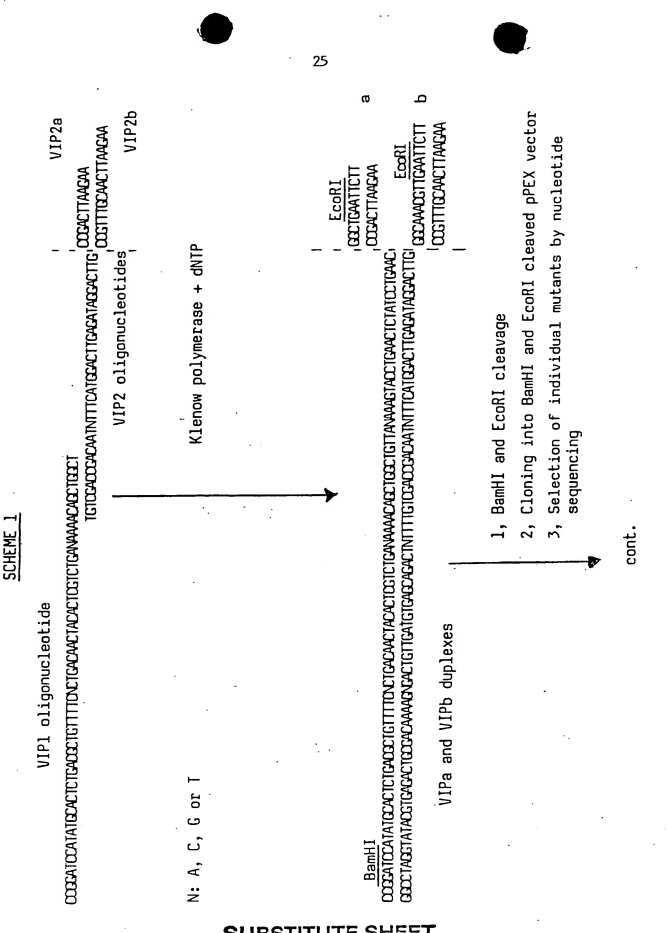
4) Biological activity of VIPa

The biological activity of recombinant VIPa was also examined in the <u>in vivo</u> assay on stimulation of bicarbonate secretion from cat pancreas according to Mutt, V. and Söderberg, U. (1959) Arkiv Kem. <u>15</u>, 63-68.

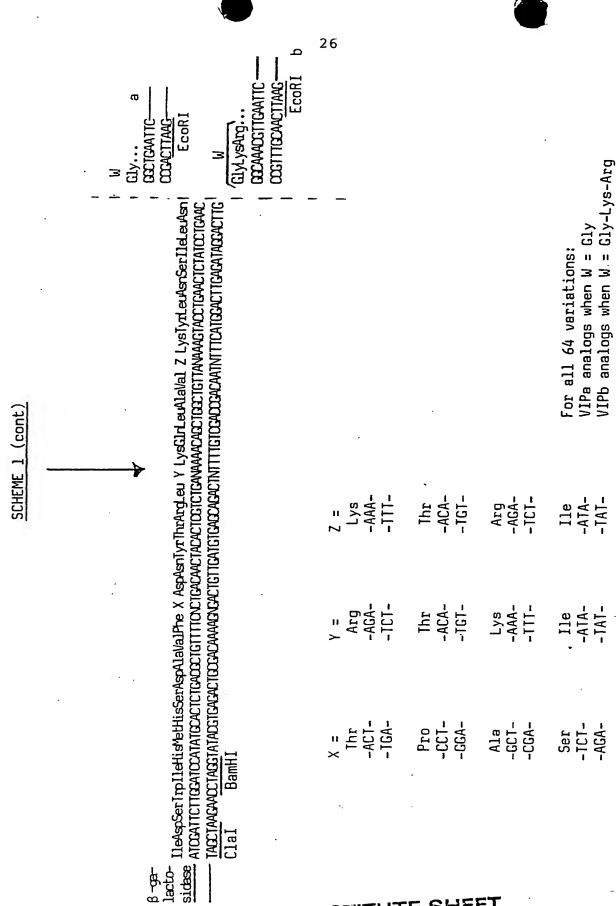
Recombinant VIPa (16 µg injected) was in this assay approximately 70% as efficient as porcine VIP.

Recombinant VIPb was, when tested at the same concentration in this assay, at least as efficient as porcine VIP.

BNSDOCID: <WO_____8905857A1_I_>



BNSDOCID: <WO_____8905857A1_1



SUBSTITUTE SHEET

Scheme 2

Sequence of the polycloning region of the pPEX vector

280 284
IleAspSerTrpIleLeu...
AICGAITCITGGAICCTCTGAGCTCTTTGGGGCCCGAGTATGCGACAGCTGGAATTCAICGAITCTTGGAICCTTTTGGAIGGTACCTTTTTGGAATTCClai BamHI Kpni Saci 8-galacto-sidase-----

SUBSTITUTE PLEAT

CLAIMS

l. A method of simultaneously producing a group of individual C-terminally extended analogs of Leu¹⁷-VIP (vasoactive intestinal polypeptide), said group consisting of equally long peptides having the amino acid sequence

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His-Ser-Asp-Ala-Val-Phe-X-Asp-Asn-Tyr-Thr-Arg-Leu-Y-1 2 3 4 5 6 7 8 9 10 11 12 13 14

-Lys-Gln-Leu-Ala-Val-Z-Lys-Tyr-Leu-Asn-Ser-Ile-Leu-10 15 16 17 18 19 20 21 22 23 24 25 26 27

-Asn-W 28 29

- wherein X represents Thr, Ser, Pro or Ala,
 Y represents Thr, Lys, Ile or Ala
 Z represents Thr, Lys, Ile or Arg, and
 W represents Gly or Gly-Lys-Arg,
- and of equally long peptides which are also Leu¹⁷-VIP-W²⁹ analogs and which arise from mutations when carrying out the method,

c h a r a c t e r i s e d by the steps of
simultaneously synthesizing a first mixture of
equally long oligonucleotides which during synthesis
are made ambiguous at the triplets coding for the
amino acid residues at positions 7 and 14 of Leu¹⁷-VIP,
and which have a few identical bases at the 3'-end.

simultaneously synthesizing a second mixture of equally long oligonucleotides which during synthesis are made ambiguous at the triplet coding for the amino acid residue at position 20 of Leu¹⁷-VIP, and which have a few identical bases at the 3'-end, these being complementary to the few bases at the 3'-end of the

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oligonucleotides in the first mixture,

mixing said first mixture with said second mixture, whereby the few bases at the 3'-end of the oligonucleotides of the first mixture anneal to the few bases at the 3'-end of the oligonucleotides of the second mixture, resulting in a mixture of partially double stranded equally long DNA sequences, which thereafter are enzymatically converted into fully double stranded DNA sequences,

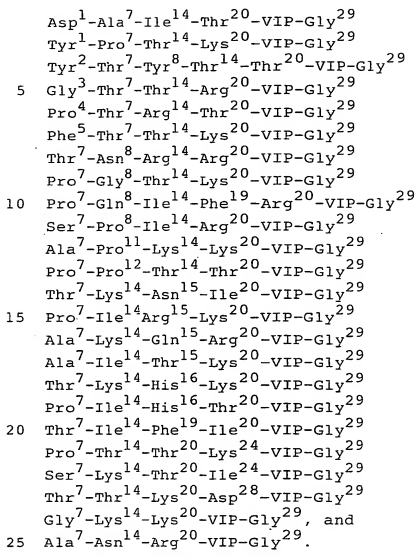
subjecting said mixture of double stranded DNA sequences, which include at the 3'-end a site for cleavage with a first enzyme and at the 5'-end a site for cleavage with a second enzyme, to cleavage, simultaneously or consecutively, with the first and the second enzyme to produce DNA sequences with 3'- and 5'-ends which are ligatable to the 5'- and 3'-ends of similarly cleaved vectors,

inserting the thus cleaved DNA sequences into the thus cleaved vectors by ligation to produce a mixture of vectors, which are then transformed into hosts in per se known manner,

propagating said hosts to form colonies, which are analyzed one by one to establish those DNA sequences which code for a single protein,

propagating separately under expression conditions those hosts which contain vectors having identified protein coding sequences, the C-terminal portions of which correspond to an individual Leu¹⁷-VIP-W²⁹ analog, whereupon the separately expressed proteins are cleaved to release the C-terminal Leu¹⁷-VIP-W²⁹ analogs forming a group of said individual equally long peptides.

2. A method according to claim 1, wherein W represents Gly and the $\text{Leu}^{17}\text{-VIP-W}^{29}$ analogs which arise from mutations when carrying out the method are



3. A method according to claim 1, wherein W represents Gly-Lys-Arg and the Leu 17 -VIP-W 29 analogs which arise from mutations when carrying out the method are

30 $Phe^{2}-Pro^{7}-Thr^{14}-Arg^{20}-VIP-Gly^{29}-Lys^{30}-Arg^{31}$ $Val^{3}-Thr^{7}-Lys^{14}-Ile^{20}-VIP-Gly^{29}-Lys^{30}-Arg^{31}$ $Pro^{4}-Thr^{7}-Thr^{14}-Lys^{20}-VIP-Gly^{29}-Lys^{30}-Arg^{31}$ $Phe^{5}-Pro^{7}-Thr^{14}-Ile^{20}-VIP-Gly^{29}-Lys^{30}-Arg^{31}$

 $_{\mathrm{Thr}^{6}-\mathrm{Ala}^{7}-\mathrm{Thr}^{14}-\mathrm{Thr}^{20}-\mathrm{VIP-Gly}^{29}-\mathrm{Lys}^{30}-\mathrm{Arg}^{31}}$ Thr 7-Val 8-Thr 14-Thr 20-VIP-Gly 29-Lys 30-Arg 31 Pro⁷-Pro¹¹-Thr¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹¹-Ile¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ 5 Thr 7-Arg 13-Lys 14-Arg 20-VIP-Gly 29-Lys 30-Arg 31 Pro⁷-Gln¹³-Lys¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹³-Thr¹⁴-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹³-Thr¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg.³¹ Ser⁷-Arg¹³-Thr¹⁴-Gln¹⁵-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Lys¹⁴-Gln¹⁵-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser⁷-Lys¹⁴-Asn¹⁵-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Thr⁷-Lys¹⁴-Asp¹⁹-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Thr7-Lys14-Gly19-Thr20-VIP-Gly29-Lys30-Arg31 Thr⁷-Thr¹⁴-Ile²⁰-Asp²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ala⁷-Ile¹⁴-Arg²⁰-Asp²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser⁷-Ile¹⁴-Thr²⁰-Lys²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Ile¹⁴-Thr²⁰-Tyr²⁵-VIP-Gly²⁹-Lys³⁰-Arg³¹

20 4. A method according to any one of claims 1-3, character is ed in that the vectors are plasmids and the hosts are E. coli.

Ser⁷-Thr¹⁴-Ile²⁰-Val²⁶-VIP-Gly²⁹-Lys³⁰-Arg³¹.

5. A C-terminally extended analog of Leu¹⁷-VIP character is ed in that it is chosen from the group consisting of peptides having the amino acid sequence

His-Ser-Asp-Ala-Val-Phe-X-Asp-Asn-Tyr-Thr-Arg-Leu-Y-1 2 3 4 5 6 7 8 9 10 11 12 13 14

30 -Lys-Gln-Leu-Ala-Val-Z-Lys-Tyr-Leu-Asn-Ser-Ile-Leu-15 16 17 18 19 20 21 22 23 24 25 26 27

-Asn-W 35 28 29



wherein X represents Thr, Ser, Pro or Ala, Y represents Thr, Lys, Ile or Ala Z represents Thr, Lys, Ile or Arg, and W represents Gly or Gly-Lys-Arg,

5 and

 $Asp^1-Ala^7-Ile^{14}-Thr^{20}-VIP-Gly^{29}$ Tyr¹-Pro⁷-Thr¹⁴-Lys²⁰-VIP-Gly²⁹ Tyr²-Thr⁷-Tyr⁸-Thr¹⁴-Thr²⁰-VIP-Gly²⁹ Gly³-Thr⁷-Thr¹⁴-Arg²⁰-VIP-Gly²⁹ 10 Pro⁴-Thr⁷-Arg¹⁴-Thr²⁰-VIP-Gly²⁹ Phe⁵-Thr⁷-Thr¹⁴-Lys²⁰-VIP-Gly²⁹ Thr⁷-Asn⁸-Arg¹⁴-Arg²⁰-VIP-Gly²⁹ Pro⁷-Gly⁸-Thr¹⁴-Lys²⁰-VIP-Gly²⁹ Pro⁷-Gln⁸-Ile¹⁴-Phe¹⁹-Arg²⁰-VIP-Gly²⁹ Ser⁷-Pro⁸-Ile¹⁴-Arg²⁰-VIP-Gly²⁹ Ala⁷-Pro¹¹-Lys¹⁴-Lys²⁰-VIP-Gly²⁹ Pro⁷-Pro¹²-Thr¹⁴-Thr²⁰-VIP-Gly²⁹ Thr7-Lys14-Asn15-Ile20-VIP-Gly29 Pro⁷-Ile¹⁴Arg¹⁵-Lys²⁰-VIP-Gly²⁹ 20 Ala⁷-Lys¹⁴-Gln¹⁵-Arg²⁰-VIP-Gly²⁹ Ala⁷-Ile¹⁴-Thr¹⁵-Lys²⁰-VIP-Gly²⁹ Thr⁷-Lys¹⁴-His¹⁶-Lys²⁰-VIP-Gly²⁹ Pro⁷-Ile¹⁴-His¹⁶-Thr²⁰-VIP-Gly²⁹ Thr^7 -Ile 14 -Phe 19 -Ile 20 -VIP-Gly 29 25 Pro⁷-Thr¹⁴-Thr²⁰-Lys²⁴-VIP-Gly²⁹ Ser⁷-Lys¹⁴-Thr²⁰-Ile²⁴-VIP-Gly²⁹ Thr⁷-Thr¹⁴-Lys²⁰-Asp²⁸-VIP-Gly²⁹ Gly⁷-Lys¹⁴-Lys²⁰-VIP-Gly²⁹ Ala⁷-Asn¹⁴-Arg²⁰-VIP-Gly²⁹ 30 and

Phe²-Pro⁷-Thr¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹
Val³-Thr⁷-Lys¹⁴-Ile²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹
Pro⁴-Thr⁷-Thr¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹
Phe⁵-Pro⁷-Thr¹⁴-Ile²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹
Thr⁶-Ala⁷-Thr¹⁴-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹



Thr 7-Val 8-Thr 14-Thr 20-VIP-Gly 29-Lys 30-Arg 31 Pro⁷-Pro¹¹-Thr¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹¹-Ile¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Thr 7-Arg 13-Lys 14-Arg 20-VIP-G1y 29-Lys 30-Arg 31 Pro⁷-Gln¹³-Lys¹⁴-Lys²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹³-Thr¹⁴-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Pro⁷-Pro¹³-Thr¹⁴-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser 7-Arg 13-Thr 14-Gln 15-Thr 20-VIP-Gly 29-Lys 30-Arg 31 Pro⁷-Lys¹⁴-Gln¹⁵-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser⁷-Lys¹⁴-Asn¹⁵-Arg²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Thr 7-Lys 14-Asp 19-Lys 20-VIP-Gly 29-Lys 30-Arg 31 Thr⁷-Lys¹⁴-Gly¹⁹-Thr²⁰-VIP-Gly²⁹-Lys³⁰-Arg³¹ Thr 7-Thr 14-Ile 20-Asp 24-VIP-Gly 29-Lys 30-Arg 31 Ala⁷-Ile¹⁴-Arg²⁰-Asp²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ Ser⁷-Ile¹⁴-Thr²⁰-Lys²⁴-VIP-Gly²⁹-Lys³⁰-Arg³¹ $Pro^7 - Ile^{14} - Thr^{20} - Tyr^{25} - VIP - Gly^{29} - Lys^{30} - Arg^{31}$ Ser⁷-Thr¹⁴-Ile²⁰-Val²⁶-VIP-Gly²⁹-Lys³⁰-Arg³¹

wherein all the amino acid residues are of L-confi-20 guration.

6. A plasmid containing a gene coding for a C-terminally extended analog of Leu¹⁷-VIP, c h a r a c - t e r i s e d by a DNA sequence which codes for a C-terminally extended analog of Leu¹⁷-VIP as defined in claim 5.

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INTERNATIONAL SEARCH REPORT

International Application No PCT/SE88/00696

I. CLASSIFICATION OF SUBJECT MA (if several classification symbols apply, indicate all)

According to International Patent Classification (IPC) or to both National Classification and IPC4

C 12 N 15/00, C 12 P 21/02, C 07 K 7/10, C 07 H 21/00

II. FIELDS SEARCHED

Minimum Documentation Searched 7

Classification System | Classification Symbols

IPC 4 C 07 H 21/00 - /04; C 07 K 1/00 - /02; C 07 K 7/10; C 12 N 15/00; C 12 P 21/00 - /02

US C1 : 435:68-71, 170-172.3

Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields Searched

SE, NO, DK, FI classes as above. Data base search: WPI/L,CA

Category •	Citati	on of Do	ERED TO BE RELEVANT® ocument, 11 with indication, where appropriate, of the relevant passages 12	Relevant to Claim No. 13
X Y	EΡ,	A2,	O 184 309 (BEECHAM GROUP PLC) 11 June 1986 See page 3 line 23, page 6 lines 35-38	5-6 1-4
; ; ;		&	JP, 61129198 US, 4737487	
Х	EP,	A2,	O 225 020 (BEECHAM GROUP PLC) 10 June 1987 See page 3 line 24 page 5 lines 26-28 JP, 62116595	5 – 6
X	us,	A, &	4 605 641 (DAVID R. BOLIN) 12 August 1986 See column 3, line 63, claim l US, 4734400	5-6
Y	WO,	Al, &	85/02198 (AMGEN) 23 May 1985 EP, 0150572 JP,T,61500250/	1-4

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- "&" document member of the same patent family

Date of Mailing of this International Search Report

IV. CERTIFICATION

Date of the Actual Completion of the International Search

1989-02-21

1989 -03- 1 3

International Searching Authority

Signature of Authorized Officer

Swedish Patent Office

Vonne Siösteen

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[&]quot;A" document defining the general state of the art which is not considered to be of particular relevance

III. DOCU	III. DOCUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)					
Category *	Citation of Document, with indication, where appropriate, of the relevant passages	· Relevant to Claim No				
А	EP, A1, O 194 OO6 (IMPERIAL CHEMICAL INDUS- TRIES PLC) 10 September 1986 & JP, 61275300	1-4				

Form PCT:ISA/210 (extra sheet) (January 1985)

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